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(54) Method for determining analyte employing probe with replicable RNA template

Verfahren zur Bestimmung eines Analyten unter Verwendung einer replizierbare RNS enthaltenden Sonde

Procédé pour la détermination d'un analyte utilisant une sonde qui contient une matrice d'ARN répliable

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Description

[0001] This application relates to the detection of organisms in the clinical and food environment and more particularly relates to new RNA template probe constructions used with RNA directed RNA polymerase amplification hybridization assays.

[0002] The need to detect low levels of pathogenic organisms in bodily fluids and foodstuffs is of primary importance in the diagnosis, treatment, and prevention of disease. Current approaches to detect such organisms include their isolation and culture *in vitro* or by direct detection of an analyte specific to the organism. Such analytes may be either the nucleic acid genome of the organism, a ribonucleic acid produced by transcription of its genome during propagation, or proteins or metabolites specifically made by or elicited in the host by the infecting organism.

[0003] Methods to detect the presence of specific nucleic acids in a sample are well understood and utilize the same general principle: the high-affinity interaction between one strand of nucleic acid bearing a specific sequence of nucleotides for another strand of nucleic acid possessing the complementary sequence to form a hybrid, or double-stranded structure. The formation of such hybrids can be made to be highly specific ("stringent") by adjustment of the conditions under which this hybridization takes place such that hybridization will not occur unless the sequences are precisely complementary. If total nucleic acid from the sample is immobilized on a solid support such as a nitrocellulose membrane, the presence of a specific "target" sequence in the sample can be determined by the binding of a complementary nucleic acid "probe" which bears a detectable moiety. Such moieties may be radioactive nuclei, fluorescent compounds, or ligands which are recognized by protein conjugates bearing catalytic activities capable of generating colored, luminescent, or fluorescent products. After removal of non-hybridized probe by washing the support, the amount of target is determined by the amount of detectable moiety present.

[0004] In practice, the sensitivity of such assays is limited by the number of labelled moieties which one may physically incorporate into the probe nucleic acid. In the case of radioactively-labelled probes, the practical limit is about 10^4 target molecules. However, to achieve this sensitivity requires probes of very high specific activity which disadvantageously possess a very limited useful lifetime coupled with a detection step (i.e.-autoradiography) requiring several days. Other labelling methods utilizing fluorescent, chemiluminescent, or enzymatic detection, although more rapid, usually do not exceed the sensitivity of isotopically-labelled probes. Since most organisms of clinical interest do not contain more than 50,000 copies of any nucleic acid suitable for use as a target, the utility of such methods is restricted to the detection of large numbers of organisms. The level of infectious agents in clinical specimens or foodstuffs, however, often does not exceed a few (1-10) per sample. Thus, a need exists for methods able to detect such low levels.

[0005] It is one aspect of the present invention to improve the sensitivity of nucleic acid hybridization assays.

[0006] One approach for the detection of low levels of DNA utilizes a DNA-dependent DNA polymerase to directly replicate the DNA target to increase its numbers to easily detectable levels. This approach is termed "polymerase chain reaction" (PCR). Saiki, R.K., Scharf, S., Faloona, F., Mullis, K.B., Horn, G.T., Erlich, H.A., and Arnheim, N., "Enzymatic Amplification of Beta-globin Genomic Sequences and Restriction Site Analysis for Diagnosis of Sickle Cell Anemia", *Science* 230:1350-1354 (1985); Saiki, R.K., Gelfand, D.H., Stoffel, S., Scharf, S.J., Higuchi, R., Horn, G.T., Mullis, K. B., and Erlich, H.A., "Primer-directed Enzymatic Amplification of DNA with a Thermostable DNA Polymerase", *Science* 239:487-491 (1988); Erlich, H.A., Gelfand, D.H., and Saiki, R.K., "Specific DNA Amplification", *Nature* 331:461-(1988); and Mullis *et al.*, European Patent Application Nos. 200362 and 201184 (see also U.S. Patents 4,683,195 and 4,683,202). In the PCR method, two DNA oligonucleotides are hybridized to specific, separated sites on the complementary strands of target DNA separated by heat treatment. The annealed probe(s) then are used to prime the synthesis of the complementary strand(s) with a DNA-dependent DNA polymerase to yield two double-stranded copies of the target. After synthesis, the strands in the resulting duplex DNA products are again separated by heat treatment, the oligonucleotide probes (present in vast excess) are reannealed to both the target and the replicated strands, and incubated further with the polymerase. Since both the target DNA and the DNA strands generated by the enzymatic extension of the oligonucleotide probes serve as templates for the primer-directed replication, the amount of specific product DNA doubles with each round until a large number of product strands (for example, one million copies) are generated.

[0007] In practice, this technique is limited by the requirement that the target for amplification be DNA (as opposed to RNA), and by the occurrence of false positives generated by hybridization of probes to homologous sites in non-target DNA which fortuitously generate similar replication products. Moreover, although target DNA may be detected with very high sensitivity, the numbers of targets present in the sample is difficult to determine without adding significantly to the complexity of the assay. Since the number of infectious agents is often important in evaluating the treatment protocol for disease, this amplification approach is disadvantageously limited because it provides qualitative rather than quantitative results.

[0008] It is another aspect of the present invention to provide methods and probes which may be more advantageously used with quantitative assays.

[0009] Another approach to improving the sensitivity of nucleic acid detection is to employ a hybridization probe coupled to or incorporated within an RNA molecule that may be replicated by (i.e.-is a template for) an RNA-directed RNA polymerase. For example, a number of means to generate RNA probes by derivitizing MDV-1 RNA, a 221 nucleotide RNA template for the RNA-dependent RNA polymerase, Q β replicase, are provided by Chu, B.C.F., Kramer, F. R., and Orgel, L.E., "Synthesis of an Amplifiable Reporter RNA for Bioassays", Nucl. Acids Res. 14: 5591-5603 (1986); Lizardi, P.M., Guerra, C.E., Lomeli, H., Tussie-Luna, I., and Kramer, F.R., "Exponential Amplification of Recombinant-RNA Hybridization Probes", Bio/Technology 6:1197-1203 (October, 1988); and European Patent Application 266,399 (EP Application No. 87903131.8). The replicatable RNA-probe construct may be advantageously employed in a sensitive hybridization assay, such as that described by Ranki, et al., U.S. 4, 563,419; Soderlund, G.B., U.S. 2,169,403; Stabinsky, U.S. 4,751,177; and Syvanen, et al., Nucl. Acids Res. 14:5037-5048. Following hybridization, the RNA-probe chimera associated with the target is replicated to generate amounts of RNA which may be easily detected by a variety of means (for example, by fluorescence using a dye such as ethidium bromide or propidium iodide). Since the MDV-1 RNA template for Q β replicase is doubled in number every 20 seconds *in vitro*, an exponential increase (for example, a billion-fold) in the number of RNA molecules occurs within a few minutes at a single temperature, and proceeds at that rate until the number of templates exceeds the number of active enzyme molecules in the reaction. After that point, the amount of template RNA increases linearly with time. As a consequence, in reactions given a sufficient period of time to reach this linear phase (for example 15 minutes for 100 molecules), the amount of amplified product RNA will be directly related to the logarithm of the number of template RNAs initially added (Lizardi et al., *supra*). Since the initial number of template-probes is proportional to the amount of targets, the number of targets present in the sample being examined may be advantageously quantitated over a very wide range in contrast to the so-called PCR amplification techniques.

[0010] It is yet another aspect of the present invention to take advantage of the properties of such RNA templates for the RNA dependent, RNA polymerase, Q-beta replicase.

[0011] Heretofore, there existed a limited number of means for joining the replicatable RNA to a probe. To be useful, such probe-RNA constructs must either serve directly as templates for the replicase or allow for the removal of the probe moiety prior to replication. In one approach, the probe is coupled to the RNA via a cystamine moiety containing a disulfide (-S-S-) linkage which can be cleaved prior to replication (Chu et al., *supra*). However, this method suffers from the need for several synthesis steps that increase the cost in labor of producing such probes. In addition and as will be readily recognized by those skilled in the art, such disulfide linkages are subject to premature cleavage by reducing agents (for example, glutathione) which occur naturally in many biological samples.

[0012] It is still another aspect of the present invention to avoid the limitations imposed by such disulfide linkage constructions.

[0013] In another approach, the probe sequence may be incorporated within the sequence of the replicatable RNA (Lizardi et al., *supra*). However, the foreign RNA sequence in many such constructs either disadvantageously affects the ability of the RNA to be efficiently replicated, or is spontaneously deleted during replication. For example, among a collection of eight such constructs, two were replicated faithfully, but six either replicated poorly or suffered deletion of the probe sequence during replication. Since such deletion events affect the rate of replication and occur randomly with time, the level of the RNA products obtained in the linear phase of the amplification cannot be used to assess target level. Moreover, for many target organisms, only a limited number of sequences (for example, one or two) may be used as hybridization targets because of a high degree of sequence homology between the target and nucleic acids in uninfected material. Thus, the additional requirement that the probe sequence be stably and efficiently replicated limits the general application of this approach.

[0014] It is still yet another aspect of the present invention to provide probe-template RNA constructions and methods which overcome this limitation.

[0015] Linking the probe sequence to either the 3' or 5' termini of template RNAs via the phosphodiester linkage normally found in RNAs, although simple to accomplish by a variety of means, has been reported to strongly inhibit replication. For example, ligation of a short oligoribonucleotide, A₁₀, to the 5' nucleotide of MDV-1 RNA rendered the RNA unable to replicate exponentially (Miele, Ph.D. thesis, Columbia University, 1982). Attachment of additional nucleotides to the 3' terminus of other template RNAs similarly inhibits their replication by Q β replicase. For example, addition of between 10-20 cytidylate residues to the 3' terminus of Q β phage RNA abolishes its template activity (Gillis, E., Devos, R. and Seurinck-Opsomer, C., Arch. Int. Physiol. Biochem. 84: 392-393 (1976)); addition of a short oligoadenylyate tract has a similar effect (Gilvarg, C. Jockusch, H. and Weissmann, C., Biochem. Biophys. Acta 414, 313-8 (1975); see also Devos, R., van-Emmelo, J., Seurinck-Opsomer, C., Gillis, E., and Fiers, W., Biochim. Biophys. Acta 447:319-27 (1976)). WO 87/06270 to Chu et al. intimate that attachment of an affinity molecule might be possible to the RNA template provided it is done through a purine linkage so that may be subjected to an acid depurination cleavage procedure prior to replication. Accordingly, the art has shown this technique does not work and thus this approach has not been used as a means to attach probe sequences to the replicatable RNA.

[0016] It is yet still another aspect of the present invention to provide new probes and methods for their use which

allow for greater flexibility of the placement of the nucleic acid probe sequences in conjunction with the autocatalytic RNA polymerase templates.

[0017] It is still yet another aspect of the present invention to provide replicatable RNA templates employing target specific, nucleic acid probe sequences attached to the 5' terminus of the replicative RNA.

5 [0018] It is a further aspect of the present invention to provide probes and methods for their use involving target specific, nucleic acid probes attached to the 3' terminus of a Q β replicative RNA.

[0019] Foster and Symons (Cell, 50:9-16 (1987)) and Uhlenbeck (Nature, 328:596-600 (1987)) have described RNA sequences, found in plant virusoids and elsewhere, possessing the unique property of undergoing autocatalytic cleavage in the presence of divalent cations. One such structure, now referred to as the "hammer-head" structure, involves the junction of three double-stranded helices at a region of defined and absolute sequence conservation. Uhlenbeck has shown that the "domains" of this structure may be on separate RNA molecules provided that they combine with each other through hybridization. V. Gerlach (Adeleide, Australia) has used an RNA bearing a portion of such a structure to cleave another totally unrelated RNA bearing the remaining elements, showing that in vitro, the message for chloramphenicol acetyl transferase may be targeted for cleavage by selecting appropriate RNA sequences to form the proper structure.

10 [0020] These "hammer-head" like structures have acquired the name "ribozymes", the name used herein, since one of the two RNAs in the reaction, like a true catalyst, is not consumed during the reaction and can be used to cleave unlimited numbers of the substrate RNAs.

15 [0021] As will become apparent to those skilled in the art, since as little as a single template RNA molecule can lead to detectable product, any approach in which a hybridization probe is amplified is limited by the background which will be produced by replication of probe molecules carried through the hybridization process irrespective of the presence of target. For example, in a hybridization assay employing a replicatable RNA probe in which target nucleic acid was immobilized on nitrocellulose, it was found that 10⁵ probe molecules were bound, regardless of whether target was present or not (Lizardi et al., *supra*). Such background limits the sensitivity of such assays since targets present in numbers equal to or below this level will not contribute significantly to the yield of amplified product and thus will not be reliably determined.

20 [0022] It is a yet further aspect of the present invention to provide RNA template probe construct and methods which are less sensitive to background limits.

25 [0023] One approach to reducing this background employs a method by which the target-probe complex is reversibly bound to the support ("reversible target capture"). After hybridization and immobilization, the complex is eluted from the support, which is then discarded with the non-specifically bound probe. The complex is then recaptured on fresh support. This process may be repeated several times to produce a significant reduction (for example, 10⁹-fold) in the amount of non-hybridized probe (see Collins, European Patent Application No. 87309308.2, for a detailed description of this target-cycling technique).

30 [0024] It is further still yet another aspect of the present invention to provide constructs and probes which may be used with such reversible target capture techniques.

35 [0025] In accordance with the principles and objects of the present invention there are provided novel probes employing replicatable RNA templates in conjunction with target specific probe sequences attached to either or both the 5' and 3' terminus of the RNA templates.

40 [0026] In the present invention, a pair of probes are constructed to bind adjacent sites in the target nucleic acid. One of the pair of probes preferably comprises a Q β RNA template having either a 3' or 5' extension bearing a probe sequence element separated from the replicatable RNA by a spacer sequence. The second probe ideally comprises an oligonucleotide coupled to a release agent comprising a nuclease (or an affinity agent which in turn can bind a nuclease) which directs endonucleolytic cleavage within the spacer sequence of the first probe. The nuclease is ideally any member of a class of nucleases which are inactive until a cofactor essential for its activity is added. Both probes may be advantageously employed in a hybridization assay in which the target nucleic acid along with the bound probes are immobilized. After washing away the bulk of the unbound probes, the cofactor required for the activity of the nucleases is added, causing cleavage and release of the replicatable RNA bearing a short extension into solution. This RNA may then be advantageously amplified by Q β replicase.

45 [0027] Thus, the present invention provides a method of detecting a target nucleic acid having a first target nucleic acid sequence and a second target nucleic acid sequence, said method comprising the steps of:

a. contacting a sample suspected of containing said target sequence with:

50 i. a first probe comprising a nucleic acid sequence which is substantially complementary to said first target nucleic acid sequence and is linked, through a spacer sequence to an RNA template which is replicable by an RNA directed RNA polymerase, wherein said spacer sequence comprises a ribonucleotide sequence;

ii. a second probe comprising a nucleic acid sequence which is substantially complementary to said second

target sequence and is linked to a release agent capable of cleaving said spacer sequence;

- 5 b. providing hybridization conditions that allow said first probe to hybridize to said first target nucleic acid sequence and said second probe to hybridize to said second target nucleic acid sequence;
- c. activating said release agent;
- d. allowing replication of said RNA template; and
- e. detecting RNA which is replicated from said RNA template, thereby detecting said target nucleic acid.

10 [0028] The invention also provides a method of detecting a target nucleic acid having a first target nucleic acid sequence and a second target nucleic acid sequence, said method comprising the steps of:

- a. contacting a sample suspected of containing said target sequence with:

- 15 i. a first probe comprising a nucleic acid sequence which is substantially complementary to said first target nucleic acid sequence and is linked, through a spacer sequence to an RNA template which is replicable by an RNA directed RNA polymerase, wherein said spacer sequence comprises a ribonucleotide sequence;
- ii. a second probe comprising a nucleic acid sequence which is substantially complementary to said second target sequence and is linked to a DNA oligonucleotide comprising a nucleotide sequence that is substantially complementary to a ribonucleotide sequence in said spacer sequence and which is capable of cleaving said spacer sequence on the addition of ribonuclease H;

- 20 b. providing hybridization conditions that allow said first probe to hybridize to said first target nucleic acid sequence and said second probe to hybridize to said second target nucleic acid sequence;
- c. adding ribonuclease H to cause cleavage of the spacer sequence;
- d. allowing replication of said RNA template; and

- e. detecting RNA which is replicated from said RNA template, thereby detecting said target nucleic acid.

[0029] A number of other embodiments of smart probe systems employing such extended Q β templates are provided.

30 [0030] While the most preferred probes of the present invention will employ target specific RNA sequences at both the 3' and 5' termini of the RNA replicatable template, the present invention will also work with DNA probes in these locations, however, it will be readily perceived that there will be greater difficulty in constructing and producing a mixed DNA-RNA probe of the present invention. The sequences required to form the ribozyme, however, must be RNA (and as specified in the ribozyme structure depicted in Figure 1A) in order for the ribozyme structure to form satisfactorily and be subject to inducible cleavage. Such cleavage will, in the most preferred embodiment, occur in the presence of 35 magnesium which also serves as a cofactor for Q β replicase directed replication of the RNA template, ideally a mid-variant. While manganese does work, it is less preferred since template specificity may be altered and, therefore, replication may not occur with the desired fidelity.

40 [0031] The most preferred RNA template is MDV-1 or midivariant, a 221 nucleotide RNA, described by Chu et al., WO 87/06270, and found in cells infected with bacteriophage Q β . MDV-1 is a common contaminant of Q β replicase preparations. Q β replicase acts upon (e.g. replicates) RNA templates which have particular primary and secondary structures such as MDV-1 and other structures. MDV-1, while replicated along with the Q β RNA genome, appears to have no known function. It is also known to exhibit some natural variation in the 5' and 3' termini, generally simply seen as an addition or deletion of a GC pair, shown by way of explanation with 4 GC pairs in Figures 1 and 2 and with 3 GC pairs in Figures 3 and 4.

45 [0032] The more preferred probe embodiments of the present invention will employ a relatively "short" target specific probe sequence at the 3' terminus of the RNA template. Host preferably, this target specific sequence will have a length of about 4 to about 12 bases and ideally, a length of about 5 bases. Such a relatively short length is preferred because following cleavage of the ribozyme those will favor disassociation of the replicatable RNA template from the target sequence at a typical cleavage temperature of about 55°C.

50 [0033] It will of course be readily recognized by those skilled in the art that increasing the length of the target specific probe sequence on the 5' terminus of the replicatable RNA template will generally foster increasing target specificity. It is also most advantageous that such a probe have a sufficient length such that attempted replication of the template with this 5' terminus target specific probe still attached will result in template collapse prohibiting further replication. On the other hand, if both the 5' and 3' target specific probes are short probes, e.g. only 5 nucleotides in length, then 55 in this embodiment one has the possibility of constant formation and turnover of ribozymes resulting in a yet more rapid cleavage of replicatable RNA templates. The disadvantage of this embodiment compared to the most preferred embodiment employing a 5'terminus target specific sequence of substantially greater length is that one would expect some loss in target specificity.

[0034] A preferred embodiment (shown in Figure 2) utilizes a probe construction wherein the first nucleic acid probe sequence specific for the RNA target is appended to the 5' leader sequence of the replicatable RNA template as in the preferred embodiment. Unlike the most preferred embodiment, however, the 3' terminus of the replicable RNA template does not incorporate a target specific nucleic acid probe sequence. Rather, a second probe comprising a second nucleic acid probe sequence specific for a second RNA target sequence, half of the ribozyme structure, and a third nucleic acid probe sequence specific for a portion of the 5' RNA template stem sequence is utilized. As with the most preferred embodiment, hybridization of both probes with the RNA target results in the formation of a cleavable ribozyme in which the RNA target provides a requisite 5'-GAAA-3' sequence.

[0035] The most preferred methods of the present invention comprise contacting the preferred probe constructions of the present invention with single-stranded target nucleic acid under conditions allowing hybridization of the first and second nucleic acid probe sequences with the first and second target sequences. "short" second nucleic acid probe sequence at the 3' end of the RNA template will be of insufficient length to stably hybridize to the RNA target. The released RNA template may then be advantageously replicated by Q β replicase and the replicated RNA template detected as an indication of the presence or absence of the target nucleic acid. The most ideal methods will include a separation step prior to cleavage induction whereby non-hybridized probe is removed and therefore unable to participate as an available template in the amplification cycle by Q β replicase. Removal may be accomplished by a variety of means such as for example by immobilizing the probe - target hybridization complex via another insolubilized target specific probe, itself directly or indirectly immobilized upon a solid surface such as a capture bead, dipstick, etc. Such so called "sandwich" techniques are well known by those skilled in the art and need not be explored in detail here.

Brief Description of the Drawings

[0036] Further understanding of the principles, objects and other aspects of the invention may be had by study of the accompanying drawings wherein:

Figure 1A shows hybridization of a midivariant with an RNA target forming the ribozyme (the "circular" structure contained within the box) showing the cleavage site;

Figure 1B illustrates the resultant cleavage products obtained by the addition of divalent cation to the probe-target complex of Figure 1A;

Figure 2 illustrates an alternative embodiment wherein a first probe comprising an RNA template and a 5' terminus target specific sequence hybridizes to the RNA target and wherein a second probe sequence hybridizes to both the RNA target and a 5' stem sequence of the RNA template of the first probe whereby a ribozyme is formed;

Figure 3 illustrates graphically the general construction scheme of a cDNA clone for use in generating probes of the instant invention;

Figures 4A and 4B illustrate graphically the use of the cDNA clone produced in Figure 3 in one construction of a vector to produce the most preferred probe of the instant invention; and

Figures 5A and 5B graphically depict the most preferred embodiment employing a pair of probes, one of which comprises the midivariant-1 attached to a target specific probe through a cleavable spacer element.

[0037] In an examination of the results of Miele (supra) and others, a set of recombinant MDV-1 RNAs bearing 3' terminal extensions of various lengths was generated by transcription of a cDNA clone described by Lizardi et al. (supra) which had been digested variously with restriction enzymes whose cleavage sites occur downstream of the MDV cDNA. It was surprisingly discovered that, contrary to the general teachings of the literature, a 3' appendage as long as 183 nucleotides in length to the 3' terminus of the recombinant RNA template allowed sufficient replication to occur such that amplified RNA products could be detected when as few as 10⁴ molecules bearing such 3' extensions were used to initiate a Q β replicase reaction. The amplified product RNAs were identical in size to the RNA lacking a 3' extension, indicating the 3' extension was not replicated with the template portion of the molecule. This surprising and totally unexpected result has significant implications for the application of replicatable RNAs as probes in amplified hybridization assays since it provides for greater flexibility in the placement of probe sequences on the replicatable RNA.

Comparative Example 1: Production of a Ribozyme: Q β template RNA probe Chimera.

[0038] The RNA construct illustrated in Figure 1 is produced by the following method. Two synthetic oligonucleotide

primers are produced by methods familiar to those skilled in the art. The first contains one strand of a number of 5
recognition sequences of the restriction endonucleases used for subsequent constructions (see Figure 4A), joined to a portion 10
complementary to the sequence proximal to but not containing the ultimate 3' nucleotide, A, of the plus strand 15
of midivariant RNA, a template for the enzyme Q β replicase. As will be understood by those skilled in the art, the exact 20
length of a sequence complementary to this region of the RNA must be of sufficient length to form an intermolecular 25
hybrid and displace the intramolecular base pairing present in the midivariant RNA in this region. The second oligo- 30
nucleotide contains, in order from its 5' to 3' end, the sequence comprising one strand of the promoter for a DNA- 35
dependent RNA polymerase read in the same sense as the direction of transcription. In this case, said promoter is that 40
for the RNA polymerase encoded by the bacteriophage T7, a highly efficient enzyme, although as will be understood 45
by those skilled in the art, promoters for other RNA polymerases can be used as well. This sequence element is 50
followed, in the 3'direction by a cleavage site for a restriction endonuclease which produces a staggered break in 55
double-stranded DNA, and which leaves a protruding single-stranded 3' terminal extension. In this example, this en- 60
zyme is Apa I, and its sequence is made to partially overlap the promoter for T7 RNA polymerase. This site is followed 65
by a recognition sequence for a restriction endonuclease which may be any member of a class of enzymes which 70
recognize a sequence and cleave the DNA in a region 3' to that sequence element, and which leave a staggered 75
cleavage and a protruding single-stranded 5' terminal extension. This recognition sequence is placed in such a manner 80
as to direct the cleavage of the final DNA vector construct in one strand immediately prior to the sequence encoding 85
the midivariant RNA. In this example (see Figure. 4A), this is the site for the restriction endonuclease Ear I which 90
recognizes the sequence CTCTTCN, where N is any nucleotide, and cleaves the DNA in the same strand after the 95
nucleotide N, and three nucleotides further in the complementary strand. Finally, this sequence is followed by another 100
sequence homologous to the sequence proximal to and including the 5' terminus of midivariant RNA (MDV-1 RNA) 105
which is of sufficient length to anneal and prime the synthesis of a second strand product as described below, except 110
that deoxyribonucleotides replace the ribonucleotides found in MDV-1.

[0039] The first oligonucleotide is used as a primer for the synthesis of a cDNA of midivariant RNA as shown in 115
Figure 3. MDV-1 is obtained from a Q β replicase reaction initiated without exogenous template RNA. This will generate 120
abundant levels of MDV-1 RNA because all preparations of Q β replicase holoenzyme prepared by any of a number of 125
methods typically are contaminated by low levels of this RNA. One picomole each of the first oligonucleotide described 130
above and MDV-1 RNA are mixed and annealed by heating and allowing the mixture to cool slowly. This primer-template 135
combination is used to initiate a reverse transcriptase reaction in a manner such as those described by Maniatis *et al.*, 140
A Cloning Manual. After completion of the first strand synthesis, the midivariant RNA is destroyed by mild alkali treat- 145
ment or alternately, RNase H. The second complementary strand is generated by annealing the product of this reaction 150
with the second oligonucleotide described above and extending with a DNA-dependent DNA polymerase. This product 155
can then be cloned into a selectable, replicatable DNA vector in a manner such as that described by Maniatis *et al.* 160
Potential clones are screened for the presence of each of the restriction endonuclease sites included in the oligonu- 165
cleotides by conventional methods.

[0040] After propagation in and purification from bacteria, the cloned cDNA is cleaved with the restriction endonu- 170
clease Apa I and Kpn I. The smaller fragment is purified electrophoretically, and cleaved with the restriction endonu- 175
clease Hinf I. This enzyme cleaves the cDNA at the site corresponding to position 65 in MDV-1. In the naturally occurring 180
population of MDV-1 RNAs, however, this site occurs in only a fraction of the molecules. Thus, a number of clones 185
must be screened for those which possess the appropriate sequence. Once thusly obtained, the Hinf subfragments 190
are ligated to the oligonucleotide 5'-pGA₁₀-3' annealed to the oligonucleotide 5'-pCT₁₀-3'. This ligation mixture is di- 195
gested with Apa I and Kpn I. The resultant mixture is ligated into the large fragment resulting from digestion of the 200
vector produced above with Apa I and Kpn I. This will generate a clone in which the DNA sequence encoding the 205
midivariant is interrupted by tract of A residues. Any of a number of sequences may be substituted for the A₁₀ tract 210
utilized in this example. However, A₁₀ is known to be maintained during replication of midivariant RNA, as shown by 215
Miele *et al.* (J. Mol. Biol. (1983) 281-295).

[0041] After propagation in and purification from bacteria, the cloned DNA from the construction above is cleaved in 220
two separate reactions (see Figure 4A). In the first of these, the segment bearing the complete midivariant RNA cDNA 225
is excised with the enzymes Ear I and Sma I. This digest is treated with T4 DNA polymerase in the presence of thymidine 230
triphosphate to remove three nucleotides from the 3' end of the strand bearing the sequence in the same sense as the 235
MDV-1 plus strand (e.g. the strand shown in Figure 1, its complement is the minus strand, also a replicatable RNA 240
template). The resulting modified fragment is purified by electrophoresis through polyacrylamide gels. The second 245
restriction digest is with the enzymes Apa I and Kpn I. The larger of the two fragments is purified by agarose gel 250
electrophoresis.

[0042] Two oligonucleotides are synthesized. The first of these advantageously contains the sequence 5'-CCCCT- 255
GANGA-3' followed by at least four nucleotides complementary to the sequence in the target RNA 5' to the 5'-GAAA- 260
3' element and terminating in the sequence 5'-GATC-3'. The second oligonucleotide ideally contains the sequence 265
5'-CCCGA-3' followed by at least 4 nucleotides of the sequence 3' to the 5'-GAAA-3' element in the target, except that 270

deoxyribonucleotides replace the ribonucleotides of the target. This element is advantageously followed immediately by the element 5'-GGGG-3'. Each of the oligonucleotides is phosphorylated on its 5' terminus by T4 polynucleotide kinase.

[0043] Both of the oligonucleotides are ligated to the large fragment produced by the Kpn I, Apa I digest of vector DNA in a reaction containing T4 DNA ligase and ATP such as those described by Maniatis et al (ibid). The ligation product (see Figure 4B) is purified away from unligated oligonucleotides by gel filtration. The ligation product is then ligated to the small Ear I, Sma I fragment bearing the modified terminus. The resulting "gapped" molecule is rendered fully double-stranded by treatment with a DNA-dependent, DNA polymerase. This product may be advantageously introduced into and propagated in bacteria.

[0044] As will be obvious to those skilled in the art, this series of manipulations will generate DNA clones in which the MDV cDNA portion is inserted in both its normal and inverted orientation relative to the direction of transcription by T7 RNA polymerase. The clones containing the MDV cDNA in plus strand orientation (oriented such that transcription produces an RNA containing the plus strand of MDV-1) are determined by screening for restriction endonuclease fragments of the appropriate size which are known to cleave within the MDV-1 cDNA sequence.

[0045] The DNA prepared from a clone with the MDV-1 in the plus strand orientation is cleaved at a restriction site distal to the second probe element relative to the promoter. In this example, this is the restriction endonuclease Dra I. This digested DNA is advantageously transcribed by T7 RNA polymerase *in vitro* under conditions such as those described by Milligan et al. (Nucl. Acid. Res. (1987) 15:8783-8798) to generate the RNA probe.

20 Example 2: Assembly of a Bimolecular Ribozyme Probe Set

[0046] The cDNA clone as described in Example 1 is cleaved with the enzymes Apa I and Ear I, and the large fragment is ideally purified away from the small fragment described above. An oligonucleotide having the sequence 5'-CCCGA-3' followed by 4-50 nucleotides identical to the sequence immediately 3' to a 5'-GAAA-3' element in the target, in turn followed by the sequence 5'-GGCC-3' is synthesized by any of a number of methods familiar to those skilled in the art. This oligonucleotide is annealed and ligated to the large restriction fragment, and the resulting single stranded "gap" region rendered double-stranded by the action of a DNA-dependent DNA polymerase. This DNA is introduced into and propagated within bacteria. After purification from bacteria, it may be cleaved with the restriction endonuclease Sma I, and then used as a template in an *in vitro* transcription reaction utilizing bacteriophage T7 RNA polymerase.

[0047] The second sequence probe may be generated by constructing a synthetic DNA template for T7 RNA polymerase as described by Milligan et al. (Nucl. Acid. Res. (1987)15:8783-8798), one strand of this template starting at its 5' end with at least 4 nucleotides of the sequence from the region 5' to the 5'-GAAA-3' element in the target except that deoxyribonucleotides replace the ribonucleotides found in the target, followed by the sequence 5'-TCNTCAG-GGGGCCCTATAGTGAGTCGTATTA-3' where N indicates any nucleotide. The other strand of the template is an oligonucleotide having the sequence 5'-TAATACGACTCACTATAG-3'. These two oligonucleotides preferably are mixed and transcribed *in vitro* as described by Milligan et al. (ibid). The product may be readily purified by any of a number of methods familiar to those acquainted with the art.

40 Example 3: Replication of modified MDV-1 RNAs bearing 3'-terminal extensions.

[0048] A cDNA clone of MDV-1 similar to that described in example 1, was obtained from F.R. Kramer of Public Health Research Institute, New York City, and contained a promoter for T7 RNA polymerase, and a modified MDV-1 cDNA sequence in which the sequence element 5'- CTCTAGATCTGAGA CTAACATAGGTCTTAACTTGACTAA-CATCGAGGCCTGCTAGAG-3' replaced the 3-nucleotide segment encoding nucleotides 64-66 in the naturally-occurring MDV-1 RNA, followed by the sequence 5'-GGGAATTC-3'. This entire sequence was cloned between the Hind III site and the EcoRI site of pSP64 (Melton et al., Nucl. Acid. Res., Vol. 12, 7035-7056). This plasmid was cleaved separately with the restriction endonucleases Sma I, EcoRI, Alu I, and Pvu II, and each of the cleaved preparations transcribed *in vitro* by T7 RNA polymerase under conditions described by Milligan et al. The RNAs produced in these reactions were purified by denaturing polyacrylamide gel electrophoresis. This procedure resulted in the generation of modified MDV-1 molecules bearing either no extension beyond the 3' terminus found in the naturally-occurring MDV-1 RNA, a 7-nucleotide 3' extension, a 24-nucleotide 3' extension, or a 183-nucleotide 3' extension.

[0049] The purified RNAs were serially diluted and 104 molecules used to prime a Q β replicase reaction as described by Chu et al. (Nucl. Acid. Res. (1986) 14:5591-603), except that the Q β replicase was that purified from the expression clone of the replicase described by Biebricher et al. (Nature (1986) 321:89-91) and physically obtained from C. Biebricher (Max Plank Institute). After 30 minutes, two microliters of each reaction was mixed with 18 microliters 95% formamide, heated to 100°C for 5 minutes, and resolved by electrophoresis on an 8% polyacrylamide gel containing 8.3M urea. The replicated recombinant RNAs were distinguished from the MDV-1 RNA which contaminates the enzyme

preparation and is also replicated, by their larger size resulting from the larger sequence replacing nucleotides 64-66, which in turn results in a lower eletrophoretic mobility. The recombinant RNA product represented approximately 50%, 86%, 72%, and 15% of the total RNA when the reaction was initiated with the modified MDV-1 RNAs bearing either no 3' extension, the 7-nucleotide extension, the 24-nucleotide extension, or the 183-nucleotide extension, respectively.

5 The recombinant RNAs produced in each case vere the same size as the recombinant RNA which lacked a 3' terminal extension, indicating that the 3' extension itself is not replicated with the rest of the molecule. This experiment has been repeated with a number of 3' terminal extensions of different sequence with comparable results.

Comparative Example 4: An assay for target nucleic acid utilizing a ribozyme probe.

10 [0050] Two deoxyribo-oligonucleotides are synthesized with the following sequence: 5'-CCCGAGGATCACCAAGCAATATTCAAAGTAGCATGACAAAAATCTGGCC-3', and 5'-CCCCCTTGACGACATC-CCGATC-3'. These are cloned into the cDNA vector described in Example 1, and the purified cloned DNA transcribed with T7 RNA polymerase after cleavage with the restriction endonuclease Dra I. The RNA product of the correct size is purified by denaturing polyacrylamide gel electrophoresis.

15 [0051] One picomole (10^{-12} mol) of this RNA is hybridized to whole human blood from patients infected with human immunodeficiency virus type 1 (HIV-1), normal human blood, or a sample to be assayed for the presence of HIV-1, in a sandwich hybridization assay such as described by Ranki et al. (ibid) in the presence of 2.5M guanidine thiocyanate and one picomole of a synthetic deoxyribo-oligonucleotide having the sequence 5'-GGAAGCACATTGACTGATATCTAACCTGGTGGTCTCATA₁₅₀-3'. With general reference to Figure 1A the hybrid is bound to a solid support by hybridization of the dA150 tract in the oligonucleotide to immobilized polydeoxythymidine such as in the well of a polystyrene microtiter plate (or a capture bead as shown in Figure 1A) coated with dT₃₀₀₀ according to the methods of Collins et al., (EP-A-0265244 and EP-A-0328829).

20 [0052] The plate is incubated at 37°C for 30 minutes after which the contents, containing unbound probes, are aspirated and discarded, and the well repeatedly washed with a buffer containing 0.5M uanidine thiocyanate. The well is then washed with a buffer containing 90mM Tris HCl (pH 7.5), and 50μl of the same buffer except containing 14mM MgC12 introduced into the well and incubated at 50°C for 15 minutes to induce ribozyme cleavage. The contents of the well containing the cleaved and released probe (see Figure 1B) are transferred to another well for replication. Five microliters of a solution containing 4mM each ATP, GTP, CTP, and UTP, 10μCi of a-³²P-CTP and 1μg of Qβ replicase are added and the plate incubated at 37°C for 25 minutes. Two microliters of the well contents are removed and added to 18μl of 95% formamide, 0.05% xylene cyanol, 0.05% bromphenol blue for halting replication. Five microliters of this are applied to the well of a denaturing 8% polyacrylamide gel, and the replicated RNAs resolved by electrophoresis until the xylene cyanol dye has migrated the length of the gel. An X-ray film is exposed to the gel for three hours and is then developed as the final step of detection. The larger RNA product of length equal to the number of nucleotides

30 between the sequence corresponding to the unappended 5' terminus of the naturally-occurring MDV-1 RNA and the unappended 3' terminus of MDV-1 RNA and including the length of the sequence element inserted into the Hinf site such as described in Example 1 (in that case 10 nucleotides), indicates the presence of HIV-1 virus or its messenger RNA in the blood sample. It will also be readily recognized that by carefully timing the replication phase and comparing the results against a suitable control, correlation can be made between the amount of replicated RNA with the amount

35 of target RNA in the original sample thereby providing quantitation results.

40 [0053] As will be readily understood, other methods of detecting the replicated RNA may be employed including immunological methods utilizing antibodies specific for RNA-DNA hybrids (e.g. requires addition of DNA oligonucleotides specific for replicated RNA sequence) or alternatively, using ethidium bromide, the fluorescence of which is enhanced by the presence of RNA polymers.

45 [0054] Example 5: A pair of probes against HIV-1, the causative agent of human acquired immune deficiency syndrome, were generated. Two pairs of partially complementary oligonucleotides were generated with the sequences:

50 (1A) 5'-GTGTGTGTGTAAGATGTTCAGCCTGATCTTACCTGTCCTATAATTTCG-3' ;

55 (1B) 5'-AATTCGAAATTATAGGACAGGTAAAGAGATCAGGCTGAACATCTTACACACACAC-3' ;

(2A) 5'-GGGCGGTGGCGAAAAAGATGTTCAGCCTGATCTCTTACCTGTCCTATAATTCG-3' ;

5 and

(2B) 5'-AATTGAAAATTATAGGACAGGTAAGAGATCAGGCTGAACATCTTTCGCGACCGCCC-3' .

10 [0055] Each of these oligonucleotide pairs.(1A and 1B; 2A and 2B) were annealed and separately cloned between the EcoRI site and the SmaI site of the plasmid used in the above experiment. After cloning, the purified plasmid DNAs were restricted with EcoRI and transcribed with T7 RNA polymerase to generate two RNAs bearing 3' extensions which were in part complementary to the same sequence in HIV-1. In the RNA produced by the clone obtained with the first 15 pair of oligonucleotides, the sequence of the replicatable RNA and the probe sequence was separated by the spacer sequence element: 5'-GUGUGUGUGU-3'. In the second clone, a spacer of the sequence 5'-GGGCG-GUCGCGCGAAA-3' was generated. Each of the RNAs was serially diluted and aliquots used to initiate Q β replicase reactions containing 1 μ g of Q β replicase

20 [0056] 90mM Tris HCl, 14mM MgCl₂ and 400 μ M each ATP, GTP, CTP and UTP. After 30 minutes at 37°C, the reactions were stopped by addition of EDTA to 20mM and ethidium bromide to 1 μ g/ml. Fluorescence of the ethidium bromide-RNA complexes was observed over an ultraviolet transilluminator. Amplified product RNA was observed in reactions initiated with at least 10⁴ molecules generated by the first clone, and 10² molecules generated by the second clone. These results indicate the sequence region immediately adjacent to the 3' end of the replicatable RNA sequence (i.e.-the spacer) strongly affects the sensitivity of probe detection. Subsequent experiments indicated that increasing 25 the reaction time did not affect the ultimate dilution yielding a detectable product by fluorescence.

25 [0057] These results provide practical guidance for using replicatable RNAs bearing 3' sequence extensions as probes for the sensitive detection of nucleic acids. By appropriate selection of the spacer sequence, RNA probes may be advantageously generated which have intrinsically greater sensitivity in the detection of target nucleic acid. Conversely, for some target nucleic acids which are present in an infectious agent at high levels (i.e.- ribosomal RNA, 30 present at up to 50,000 copies per organism), a spacer region conferring relatively poor limits of detection may be utilized to advantageously avoid the background otherwise generated by the amplification of lower levels of non-specifically bound probes. For example, if reduced in number to levels at least an order of magnitude below the limit of detection using background reduction methods such as Collins (supra), non-specifically bound probes will be incapable 35 of producing a detectable signal. Thus, the probes of the present invention comprising appropriate spacer sequences may be used advantageously to reduce the cost, complexity, and frequency of false-positive reactions in such assays.

35 [0058] As will now be recognized, a number of means may be utilized to generate such constructs including the use of T4 RNA ligase (Stefano, supra) and transcription of cDNA clones as described in the above example. It will be appreciated that the use of T4 RNA ligase advantageously allows the generation of constructs in which the probe extension is either RNA or DNA.

40 [0059] Since Q β replicase initiates synthesis of the daughter strand at the 3' end of MDV-1 RNA template and generally continues to copy the template in a 3' to 5' direction (of the template) until the 5' terminus is reached, template molecules bearing 5' extensions will generate daughter strand products (e.g. a minus strand of template) bearing 3' terminal extensions, which in turn will be replicated as observed above (e.g. the Q-beta replicase will ignore the 3' sequences appended to the end). As will be apparent to those skilled in the art, such probes may also be employed 45 in hybridization assays. As will now be additionally apparent, the effect of the spacer sequence may similarly be employed to advantageously adjust the sensitivity of such probes.

45 [0060] An important advantage issues from the present discovery which has additional application for smart probe systems which employ the target-directed cleavage of 5' or 3' sequence extensions, since such methods need not provide for the precise and complete removal of those sequences. For example and as set forth previously, a probe 50 may be generated bearing both 3' and 5' extensions whose hybridization to target produces a ribozyme structure which directs cleavage of the 5' extension from the RNA. A short 3' extension remains on the molecule.

55 [0061] A most preferred embodiment of the present invention is diagrammed in the Figures 5A and 5B wherein a probe molecule bearing an extension is hybridized to target in solution. Shortly before, after, or simultaneously therewith, a second oligonucleotide probe is hybridized to the target adjacent to the first probe. The second oligonucleotide probe is ideally coupled to a nuclease which requires a cofactor (for example, a divalent cation) for its activity. Following hybridization, the complex is captured on a solid support in a manner such as that described by Ranki *et al.* (supra), Soderlund (supra), Stabinsky (supra), and/or Syvanen (supra). After separating, such as by washing away, the bulk of non-specifically bound probes, the cofactor for nuclease activity is added. The nuclease, coupled to the terminus of

the probe proximal to the first probe when hybridized to its target, cleaves the spacer on the first probe, thereby releasing a replicatable moiety into solution. As previously indicated, the spacer sequence may be advantageously selected to maximally inhibit replication until cleavage occurs. Other variations of this approach will become apparent. For example, such probe pairs may include a replicatable RNA instead bearing a 5' extension and a nuclease-coupled probe in which the ribozyme sequence is present 5' to the probe sequence element.

5 [0062] Any of a number of nucleases may be advantageously employed as the cleavage agent. For example, micrococcal nuclease may be used since it does not cleave MDV-1 RNA (Hill & Blumenthal (1983) *Nature* 301, 350-352). Corey & Schultz (1987) *Science* 238:1401-1403, teach the construction of such oligonucleotide-micrococcal nuclease conjugates. In the most preferred embodiment, any one of a number of ribozymes with endonucleolytic cleavage activity, 10 such as those described by Haselhoff & Gerlach (*Nature* 334: 585-591 (1988)), Uhlenbeck (1987) *Nature* 328, 596-600, Cech (European Pat. Appl. WO 88/04300, June 16, 1988), Sharpen *et al.* (1989) *J. Virol.* 63, 1428-1430; or Hampel & Tritz (1988) *J. Cell. Biochem., Suppl.* 12D, Abst. #N212, p.31, is employed. Ribozymes possess a significant advantage in that the second probe bearing the ribozyme may be efficiently produced in a single step by transcription of a 15 DNA oligonucleotide of appropriate sequence such as is described by Mulligan (*supra*), thus reducing the cost and labor required to generate such reagents. Moreover, these preferred reagents advantageously produce specific cleavage events, leading to a product RNA with defined replication properties.

10 [0063] In the specific case where the target is RNA, the cleavage agent may be a small DNA oligonucleotide (for example, six nucleotides) complementary to a portion of the spacer sequence on the first probe. In this case, the cleavage is ideally effected by the addition of RNase H (which acts to cleave RNA in RNA:DNA heteroduplexes) to the 20 solution in contact with the support bearing the complex. Naturally, the means for capturing the RNA target should ideally avoid generation of such heteroduplexes in order for the cleavage event to be specific. For example, a biotinylated RNA complementary to another portion of the target RNA may be conveniently captured upon an immobilized streptavidin support.

25 [0064] In the specific case where the target is DNA, digestion of the sequence extension of hybrids with target may be effected directly by the addition of RNase H, without the requirement for a second probe bearing such an oligonucleotide.

20 [0065] Further understanding may be had by study of the additional detailed examples which follow.

30 **Example 6: Detection of *Chlamydia trachomatis* RNA**

[0066] Two oligonucleotides having the sequences:

35 (1) 5'-AATTCTATGTGATATCAGCTAGTTGGTGGGTAAAGGCCT-3' ;

40 and

(2) 5'-AATTAGGCCTTACCCCACCAACTAGCTGATATCACATAG-3'

45 were annealed and ligated into a MDV cDNA construct similar to that described by Lizardi (*supra*), which had been digested with EcoRI. This cDNA clone differed from those described by Lizardi *et al.* in that the internal insert encoded a binding site for the coat protein for phage R17. This cDNA was obtained from F.R Kramer, Public Health Research Institute, N.Y. The oligonucleotides were ligated in the orientation such that upon subsequent digestion with EcoRI, cleavage occurred downstream of the oligonucleotide with respect to the promoter for T7 RNA polymerase. The 50 digested plasmid was transcribed *in vitro* with T7 RNA polymerase under the conditions described by Mulligan *et al.* (*supra*), and the transcription product of correct length isolated by electrophoresis through polyacrylamide gels containing 8.3M urea.

[0067] A capture oligonucleotide having the sequence:

55 5'-TACACCGCTATAAACCGTAGGCTCATTGCAATTTC-3'

complementary to a region in the 16S ribosomal RNA of *Chlamydia* (*Palmer et al.*, *Chlamydial Infections (Oriel et al.*,

eds) Cambridge Univ. Press, 1986, pp. 89-92) was synthesized and tailed with ca 150 dA residues using terminal deoxynucleotidyl transferase (Supertechs) as described by Collins (supra).

[0068] Various numbers of formalin-fixed elementary bodies of Chlamydia trachomatis were lysed in a solution of 1.0mg/ml proteinase K (Boehringer Mannheim) and 1.6% Sarkosine (Sigma) at 65°C for 15 minutes in a final volume of 35μl. Thirty-five μl of a buffer containing 340 ng/ml of tailed capture oligonucleotide and 100 ng/ml of the transcription product probe was then added and solution phase hybridization allowed to occur for 30 minutes at 37°C. Fifty μl of a 0.06%(w/v) suspension of oligo-dT derivitized magnetic beads prepared according to Collins (supra) in 4% BSA, 10mM EDTA, 0.2% Sarkosine, 0.1M Tris-HCl pH8.0 and 0.05% bronopol was then added and incubated for an additional 5

5 minutes at 37°C to capture the target-probe hybrids on the beads.

10 [0069] Following capture, 0.2ml of a "wash solution" containing 1M guanidine thiocyanate (GuSCN, Fluka), 10mM EDTA, 0.04 M Tris-HCl pH7.8, 0.5% Sarkosine, 0.2% BSA and 0.1% antifoam (Thomas) at 37°C was added. The tubes were vortexed and then placed in a magnetic field (preferably such as that described in EP-A-317,826 entitled "Magnetic Separation Device and Methods for Use",). The beads were drawn to the side of the tubes and the liquid phase removed from the tubes by aspiration. Another 0.2ml of warm wash solution was added, the beads were resuspended by vortexing, and the beads recollected in the magnetic field. The beads were washed an additional time with another aliquot of wash solution.

15 [0070] The collected beads, freed of supernatant, were then resuspended in 50μl of a buffer containing 3.25M GuSCN, 65mM EDTA, 0.04M Tris-HCl, pH7.0, 0.5% Sarkosine, and 0.5% BSA, and incubated at 37°C for 5 minutes to release the target-MDV probe-capture probe hybrids. The magnetic beads were collected as before, and the supernatants removed and transferred to a fresh set of tubes, each containing 50μl of a fresh bead suspension to recapture the hybrids as described above. These beads were washed three times in the same manner as the first set, the hybrids released and recaptured- by a third set of beads. This set of beads was washed three times in the same manner, and additionally, three times with 0.2ml of a solution containing 0.1M KCl, 1mM EDTA, 10mM Tris, pH8.0, 0.5% NP-40.

20 [0071] The collected, washed beads were resuspended in 50μl of a buffer consisting of 10mM Tris-HCl, pH8.0, 1mM EDTA, and 0.5% NP-40, and incubated at 37°C for 5 minutes to elute the complexes. The beads were collected with a magnetic field, and the supernatants containing the eluted complexes transferred to a fresh set of tubes.

25 [0072] Ten μl of each of the bead elutes was added to a set of tubes containing 13μl of a solution of 173mM Tris-HCl, pH7.5, 27mM MgCl₂, and 0.77mM each ATP, GTP, CTP, and UTP (Pharmacia). Two μl of Qβ replicase (1.2μg) purified according to DiFrancesco (supra) was added, and the tubes incubated for 12 minutes at 37°C. Five μl of 20mM EDTA, 32μg/ml propidium iodide (Sigma) was then added to stop replication. Fifty μl of each stopped reaction was transferred to separate wells of a U-bottomed microtiter plate (Nunc). The plate was photographed over an ultraviolet transilluminator whose emission maximum was 365nm onto Polaroid type 667 film using a Wratten #29 filter. The exposure was adjusted to minimize the apparent fluorescence in a control well containing reaction buffer and propidium iodide alone. The above assay was performed with triplicate samples of each elementary body dilution.

30 [0073] The results are summarized in the table below. The number of "+" symbols reflects the relative fluorescence observed. "-" indicates the fluorescence did not exceed that of the control wells.

40	Elementary bodies	Fluorescence		
		Hybridization Reaction:		
		a	b	c
	10 ⁵	+++++	++++++	+++++
45	10 ⁴	+++	+++	+++
	10 ³	++	++	+
50	10 ²	-	++	-
	10	-	-	-
	1	-	-	-
55	0	-	-	-

[0074] As observed in the above experiment, the assay had a sensitivity of approximately 10³ elementary bodies.

Example 7: Detection of HIV-1 mRNA

5 [0075] The assay of Example 6 was repeated, except that the capture probe was substituted with three probes complementary to conserved regions in HIV-1 RNA. The detector MDV probe was as described in Example 5 and contained the spacer sequence 5'-GGCGGGUCGCGCGAAA-3'. This assay showed a sensitivity of 1000 HIV-1 RNA molecules.

Example 8: Selection of probes containing optimally-replicating spacer sequences

10 [0076] A library of template DNAs can be generated by synthesis of an oligonucleotide having in 5' to 3' order: (1) a short (for example, 40 nucleotides) sequence identical to target, (2) a short stretch (15 nucleotides) of random sequence, and (3) a sequence complementary to the 3'-most 30 nucleotides of MDV-1 RNA. The random sequence element may be generated with ease by addition of all four blocked nucleotides during synthesis of the oligonucleotide, a procedure easily accomplished with currently available automated synthesizers. A second oligonucleotide is then synthesized which is complementary to the nineteen 5'-most nucleotides of the first oligonucleotide. This is annealed to the first oligonucleotide and extended with a DNA-dependent DNA polymerase to produce a double-stranded DNA fragment. This fragment is then cleaved with the restriction enzyme Bst EII and the cleavage product ligated to a fragment of a MDV cDNA clone such as described by Lizardi (supra), previously cleaved with Bst EII. Two picomoles of the ligation product is then transcribed by T7 RNA polymerase to generate a population of RNAs containing 10¹² different spacer sequences.

15 [0077] Molecules within the population of RNAs which most efficiently initiate replication are advantageously selected by one of two methods: electrophoretic retardation of Q β template RNAs hybridized to their nascent initiated daughter strands (Mills et al. (1980) Biochem 19: 228-236); or isolation of the stable complex formed between Q β replicase and template RNAs which have directed initiation of a daughter strand. An example of the latter method is described below.

20 [0078] Two picomoles of RNAs bearing random sequence spacers is incubated with 5 picomoles (1.2 μ g) of Q β replicase for 5 minutes at 37°C in 25 μ l of 100mM Tris HCl, pH7.5, 14mM MgCl₂. Five μ l of the same buffer containing 2.4mM each GTP and ATP is added, and the incubation continued for an additional 5 minutes. Five μ l of buffer containing 40 picomoles of MDV-1 RNA is then added and the incubation continued for 2 minutes. The reaction is chilled to 0°C and applied to the top of a 2.2ml 10-30% glycerol gradient in 10mM Tris-HCl, pH7.5, 1mM MgCl₂ prechilled to 0°C. The gradient is spun at 55K RPM in a TLS-55 rotor (Beckman Instruments, Palo Alto, CA) for 6 hours at 4°C. The enzyme: RNA complex, which sediments 40% faster than free RNA, is collected, extracted with phenol, and ethanol precipitated. This population is enriched in RNAs which can be replicated efficiently by Q β replicase. To further enrich this population, a cDNA library of this population is first constructed. An oligonucleotide complementary to the probe region is used to prime synthesis of a single-stranded cDNA population using reverse transcriptase (Maniatis et al., supra). A second oligonucleotide is synthesized to contain, in 5' to 3' order, (1) a restriction enzyme sequence to allow cloning of the duplex cDNA, (2) the sequence of a promoter for T7 RNA polymerase, and (3) the sequence of the first 22 nucleotides of MDV-1 RNA. This oligonucleotide and the one used for priming synthesis of the first strand cDNA product are used in a PCR reaction to generate one picomole of duplex DNA using as a template the first strand cDNA. This product is purified by native polyacrylamide gel electrophoresis, and transcribed with T7 RNA polymerase to generate an RNA population which is subjected again to the complex formation-sedimentation protocol. This process may be repeated as many times as desired, but as will be recognized, will converge upon some limiting degree of enrichment. The degree of enrichment after each cycle of selection is determined by cloning the duplex cDNAs into plasmid vectors, and examining the sensitivity of detection of the RNAs produced by transcription of several (for example, 12) such clones chosen at random from the products of each cycle of selection.

45 [0079] The sensitivity of detection of the cloned RNAs may be determined as follows. The plasmid DNAs from each clone is purified by standard methods. Each is restricted with a restriction enzyme immediately downstream of the cDNA and transcribed by T7 RNA polymerase to produce an RNA product devoid of vector sequence. This RNA is purified either by gel electrophoresis or by capture onto oligo dT-magnetic beads as in the method described in Example 6 with a DNA oligonucleotide of the same sequence as the target tailed with dA₁₅₀, except that a secondary capture probe is omitted. Each of the purified transcripts is serially diluted in water and aliquots (10 μ l) added to Q β reactions as described in Example 6. The reactions are allowed to run 30 minutes at 37°C, stopped, and the minimum number of probes required to observe product determined by the highest dilution producing fluorescence. This process is repeated for clones obtained from each round of the selection process.

50 [0080] It will be readily appreciated that the above process may also be applied to the isolation of RNAs which replicate efficiently when hybridized to target. This is simply performed by first hybridizing the population of RNAs to a synthetic target prior to the formation of initiation complexes. As will also be recognized, members of such a population may contain molecules which replicate more efficiently when hybridized to target than when free. Such molecules will be preferred probes for use in hybridization assays since the molecules nonspecifically carried through the hybridization

process will be intrinsically less efficiently replicated.

Example 9: A smart probe system for the detection of HIV-1 RNA

5 [0081] An oligonucleotide of the sequence: 5'-AATTCTTAAAAATCATGGACAGGTAA
GAGATCAAGCTGAACATCTTGGAGGGAAACAGGACTGTCAGGGACCCCCCTCGGGGG-3' is synthesized. A
second oligonucleotide of the sequence: 5'-GTGACCCCCCGAAGGGGGTCCCTTGCTGACAGTCCCTC-
CAAGATGTTAGCTTGATCTC TTACCTGTCCTATGATTTTAAG-3' is synthesized. The two oligonucleotides are
annealed and ligated into a cDNA clone of a recombinant MDV RNA as described by Lizardi (supra) and previously
10 digested with Bst EII and EcoRI. The resultant clone is digested with Dra I and transcribed with T7 RNA polymerase.
The product RNA is purified on an 6% polyacrylamide gel containing 8.3M urea, and eluted from the gel after visuali-
zation either by autoradiography or UV shadowing. A template oligonucleotide having the sequence:
5'-TACCAGGTAATATACCAACGTGTGTTCTCTGGTTGACTTCTC
15 TGTTTGGGGGGAGACAGCAGTACAAATGGCAGTATTCCACAAATTCCTATAGTGAGTCGTATT AAT-3' for
a ribozyme-probe is synthesized. Two picomoles of this product is annealed to an equimolar amount "T7 promoter-
primer" oligonucleotide (Promega) and transcribed by T7 RNA polymerase as described by Mulligan (supra), except
that α -³²P CTP (1 Ci/mmol) is included in the reaction in order to label the product for visualization by autoradiography.
The transcription product is purified on a 10% polyacrylamide gel containing 8.3M urea and eluted. One nanogram of
each of the above RNAs is added to a HIV-1 assay as described in Example 7. After washing the beads for the final
20 time as described in Example 6, the beads are resuspended in 50 μ l of an elution buffer containing 200mM KCl, 10mM
MgC₁₂, and 0.5% NP-40. The beads are incubated in this buffer at 37°C for 10 minutes, and then recollected by placing
in a magnetic field. The liquid phase is collected and transferred to a fresh tube. The RNA present in 10 μ l of this elute
is amplified in a Q β reaction as described in Example 6.

25

Claims

1. A method of detecting a target nucleic acid having a first target nucleic acid sequence and a second target nucleic acid sequence, said method comprising the steps of:
 - a. contacting a sample suspected of containing said target nucleic acid with:
 - i. a first probe comprising a nucleic acid sequence which is substantially complementary to said first target nucleic acid sequence and is linked, through a spacer sequence, to an RNA template which is replicable by an RNA directed RNA polymerase; and
 - ii. a second probe comprising a nucleic acid sequence which is substantially complementary to said second target nucleic acid sequence and is linked to a release agent capable of cleaving said spacer sequence;
 - b. providing hybridization conditions that allow said first probe to hybridize to said first target nucleic acid sequence and said second probe to hybridize to said second target nucleic acid sequence;
 - c. activating said release agent;
 - d. allowing replication of said RNA template; and
 - e. detecting RNA which is replicated from said RNA template, thereby detecting said target nucleic acid.
- 45 2. A method of detecting a target nucleic acid having a first target nucleic acid sequence and a second target nucleic acid sequence, said method comprising the steps of:
 - a. contacting a sample suspected of containing said target nucleic acid with:
 - i. a first probe comprising a nucleic acid sequence which is substantially complementary to said first target nucleic acid sequence and is linked, through a spacer sequence, to an RNA template which is replicable by an RNA directed RNA polymerase, wherein said spacer sequence comprises a ribonucleotide sequence; and
 - ii. a second probe comprising a nucleic acid sequence which is substantially complementary to said second target nucleic acid sequence and is linked to a DNA oligonucleotide comprising a nucleotide sequence that is substantially complementary to a ribonucleotide sequence in said spacer sequence and which is capable of cleaving said spacer sequence on the addition of ribonuclease H;

5 b. providing hybridization conditions that allow said first probe to hybridize to said first target nucleic acid sequence and said second probe to hybridize to said second target nucleic acid sequence;
c. adding ribonuclease H to cause cleavage of the spacer sequence;
d. allowing replication of said RNA template; and
e. detecting RNA which is replicated from said RNA template, thereby detecting said target nucleic acid.

10 3. The method of claim 1, wherein said release agent is micrococcal nuclease.
4. The method of any preceding claim, wherein said RNA template is MDV-1.
15 5. The method of any preceding claim, wherein said RNA directed RNA polymerase is Q-beta replicase.

15 **Patentansprüche**

1. Verfahren zum Nachweis einer Ziel-Nukleinsäure mit einer ersten Ziel-Nukleinsäuresequenz und einer zweiten Ziel-Nukleinsäuresequenz, wobei bei dem Verfahren:

20 a. eine Probe, die vermutlich die Ziel-Nukleinsäure enthält, in Kontakt gebracht wird mit:
i. einer ersten Sonde, die eine Nukleinsäuresequenz umfaßt, welche im wesentlichen zu der ersten Ziel-Nukleinsäuresequenz komplementär ist und durch eine Spacer-Sequenz an eine RNA-Matrize, die durch eine RNA-gerichtete RNA-Polymerase replizierbar ist, gebunden ist,
25 ii. einer zweiten Sonde, die eine Nukleinsäuresequenz umfaßt, welche im wesentlichen komplementär zu der zweiten Ziel-Nukleinsäuresequenz ist und an ein Freisetzungsmittel gebunden ist, das die Spacer-Sequenz spalten kann,
30 b. Hybridisierungsbedingungen vorgesehen werden, welche der ersten Sonde die Hybridisierung an die erste Ziel-Nukleinsäuresequenz und der zweiten Sonde die Hybridisierung an die zweite Ziel-Nukleinsäuresequenz gestatten,
35 c. das Freisetzungsmittel aktiviert wird,
d. die RNA-Matrize replizieren gelassen wird und
e. die von der RNA-Matrize replizierte RNA nachgewiesen wird, wodurch die Ziel-Nukleinsäure nachgewiesen wird.

40 2. Verfahren zum Nachweis einer Ziel-Nukleinsäure mit einer ersten Ziel-Nukleinsäuresequenz und einer zweiten Ziel-Nukleinsäuresequenz, wobei bei dem Verfahren:

45 a. eine Probe, die vermutlich die Ziel-Nukleinsäure enthält, in Kontakt gebracht wird mit:
i. einer ersten Sonde, die eine Nukleinsäuresequenz umfaßt, welche im wesentlichen komplementär zu der ersten Ziel-Nukleinsäuresequenz ist und durch eine Spacer-Sequenz an eine RNA-Matrize, die durch eine RNA-gerichtete RNA-Polymerase replizierbar ist, gebunden ist, wobei die Spacer-Sequenz eine Ribonukleotidsequenz aufweist und
50 ii. einer zweiten Sonde, die eine Nukleinsäuresequenz umfaßt, welche im wesentlichen komplementär zu der zweiten Ziel-Nukleinsäuresequenz ist und an ein DNA-Oligonukleotid gebunden ist, das eine Nukleinsäuresequenz umfaßt, die im wesentlichen komplementär zur einer Ribonukleotidsequenz in der Spacer-Sequenz ist und die Spacer-Sequenz bei Zugabe von Ribonuklease H spalten kann,
55 b. Hybridisierungsbedingungen vorgesehen werden, welche der ersten Sonde die Hybridisierung an die erste Ziel-Nukleinsäuresequenz und der zweiten Sonde die Hybridisierung an die zweite Ziel-Nukleinsäuresequenz gestatten,

c. Ribonuklease H zur Spaltung der Spacer-Sequenz zugegeben wird,
d. die RNA-Matrize replizieren gelassen wird und
5 e. von der RNA-Matrize replizierte RNA nachgewiesen wird, wodurch die Ziel-Nukleinsäure nachgewiesen wird.

3. Verfahren nach Anspruch 1, wobei das Freisetzungsmittel Mikrococcus-Nuklease ist.

10 4. Verfahren nach einem der vorstehenden Ansprüche, wobei die RNA-Matrize MDV-1 ist.

5. Verfahren nach einem der vorstehenden Ansprüche, wobei die RNA-gerichtete RNA-Polymerase Q-beta-Replikase ist.

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Revendications

1. Procédé de détection d'un acide nucléique cible ayant une première séquence d'acide nucléique cible et une seconde séquence d'acide nucléique cible, ledit procédé comprenant les étapes de :

20 a. mettre en contact un échantillon suspecté de contenir ledit acide nucléique cible avec :

25 i. une première sonde comprenant une séquence d'acide nucléique qui est实质iellement complémentaire à ladite première séquence d'acide nucléique cible et est liée, par une séquence d'espacer, à une matrice d'ARN qui est répliable par une polymérase d'ARN dirigée par l'ARN ; et
ii. une seconde sonde comprenant une séquence d'acide nucléique qui est实质iellement complémentaire à ladite seconde séquence d'acide nucléique cible et est liée à un agent de libération capable de cliver ladite séquence d'espacer ;

30 b. fournir des conditions d'hybridation qui permettent à ladite première sonde d'hybrider à ladite première séquence d'acide nucléique cible et à ladite seconde sonde d'hybrider à ladite seconde séquence d'acide nucléique cible ;
c. activer ledit agent de libération ;
d. permettre la réplication de ladite matrice d'ARN ; et
35 e. détecter l'ARN qui est répliqué à partir de ladite matrice d'ARN, en détectant de cette façon ledit acide nucléique cible.

2. Procédé de détection d'un acide nucléique cible ayant une première séquence d'acide nucléique cible et une seconde séquence d'acide nucléique cible, ledit procédé comprenant les étapes de :

40 a. mettre en contact un échantillon suspecté de contenir ledit acide nucléique cible avec :

45 i. une première sonde comprenant une séquence d'acide nucléique qui est实质iellement complémentaire à ladite première séquence d'acide nucléique cible et est liée, par une séquence d'espacer, à une matrice d'ARN qui est répliable par une polymérase d'ARN dirigée par l'ARN, dans laquelle ladite séquence d'espacer comprend une séquence de ribonucléotide ; et
ii. une seconde sonde comprenant une séquence d'acide nucléique qui est实质iellement complémentaire à ladite seconde séquence d'acide nucléique cible et est liée à un oligonucléotide d'ADN comprenant une séquence de nucléotide qui est实质iellement complémentaire à une séquence de ribonucléotide dans ladite séquence d'espacer et qui est capable de cliver ladite séquence d'espacer lors de l'addition de ribonucléase H ;

50 b. fournir des conditions d'hybridation qui permettent à ladite première sonde d'hybrider à ladite première séquence d'acide nucléique cible et à la seconde sonde d'hybrider à ladite seconde séquence d'acide nucléique cible ;
c. ajouter la ribonucléase H pour provoquer le clivage de la séquence d'espacer ;
d. permettre la réplication de ladite matrice d'ARN ; et
55 e. détecter l'ARN qui est répliqué à partir de la matrice d'ARN, en détectant de cette façon ledit acide nucléique

cible.

3. Procédé selon la revendication 1, dans lequel ledit agent de libération est une nucléase de microcoques.
- 5 4. Procédé selon l'une quelconque des revendications précédentes, dans lequel la matrice d'ARN est MDV-1.
5. Procédé selon l'une quelconque des revendications précédentes, dans lequel ladite polymérase d'ARN dirigée par l'ARN est la Q-βéta-réplicase.

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FIGURE 1A

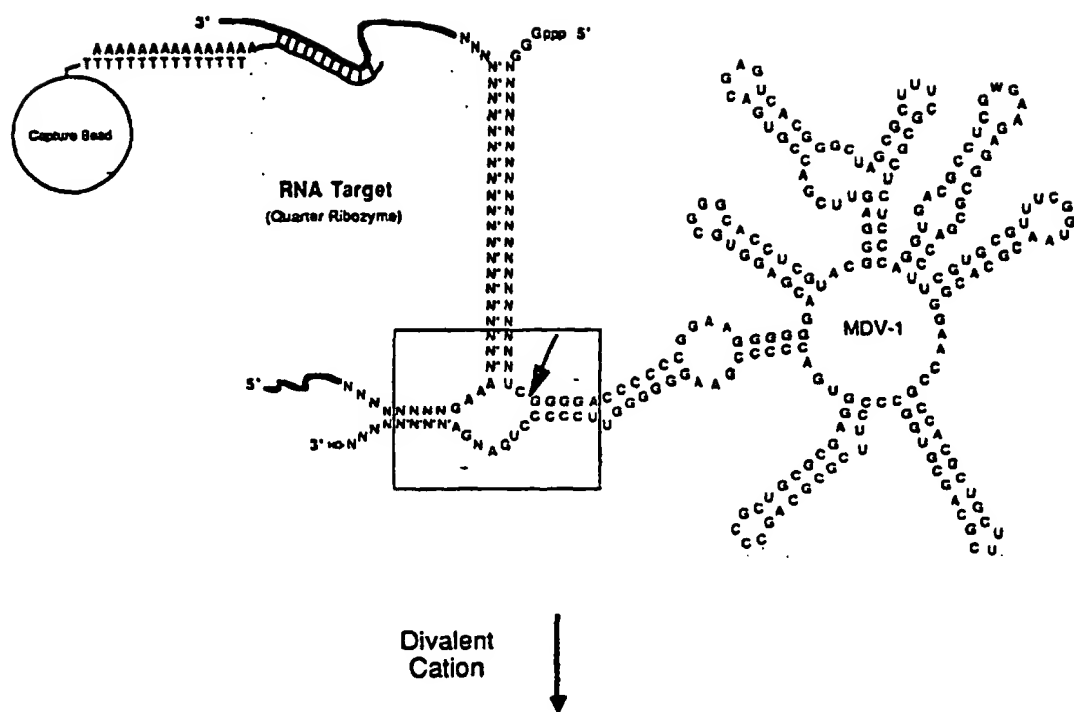


FIGURE 1B

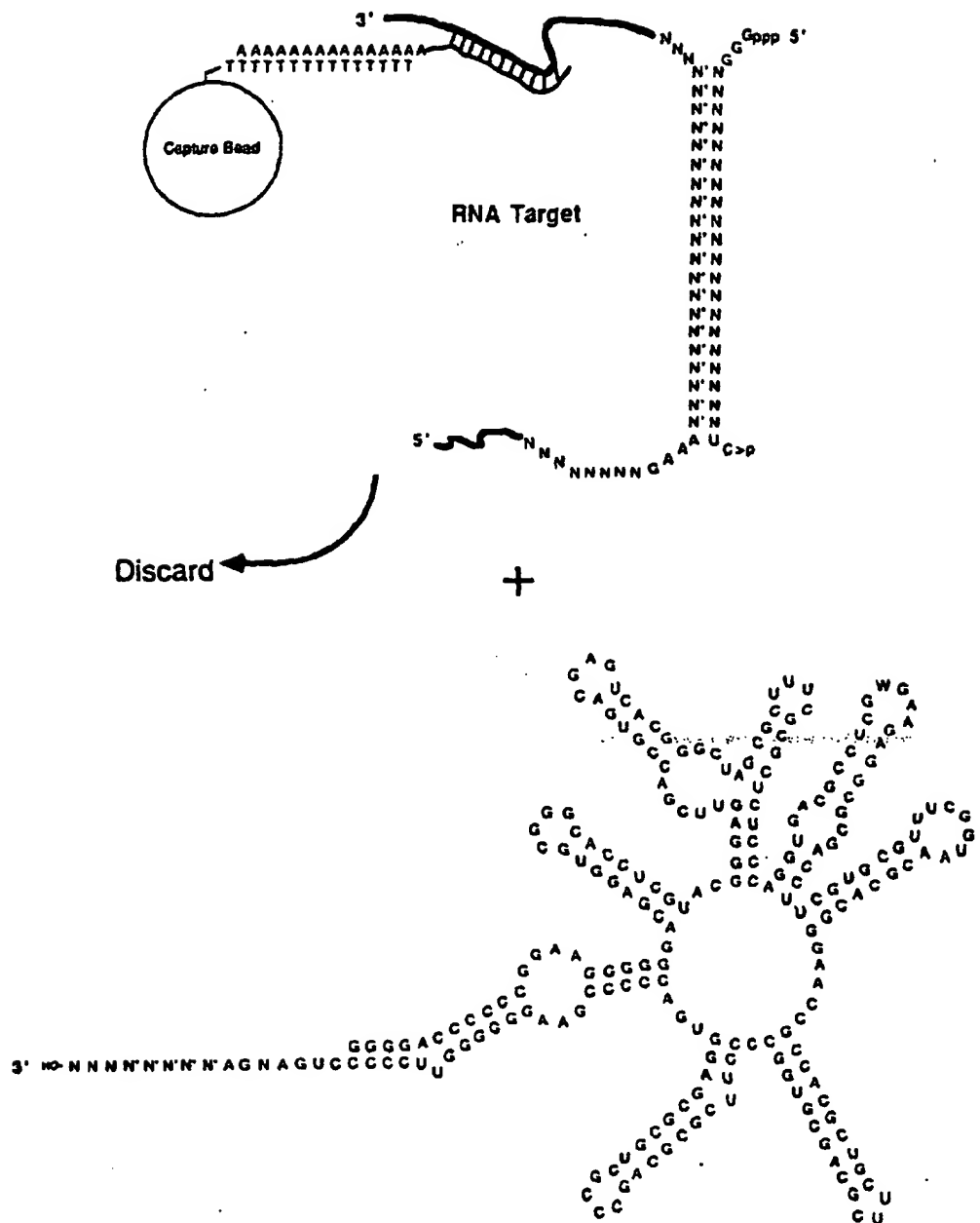
FIGURE 1B

FIGURE 2

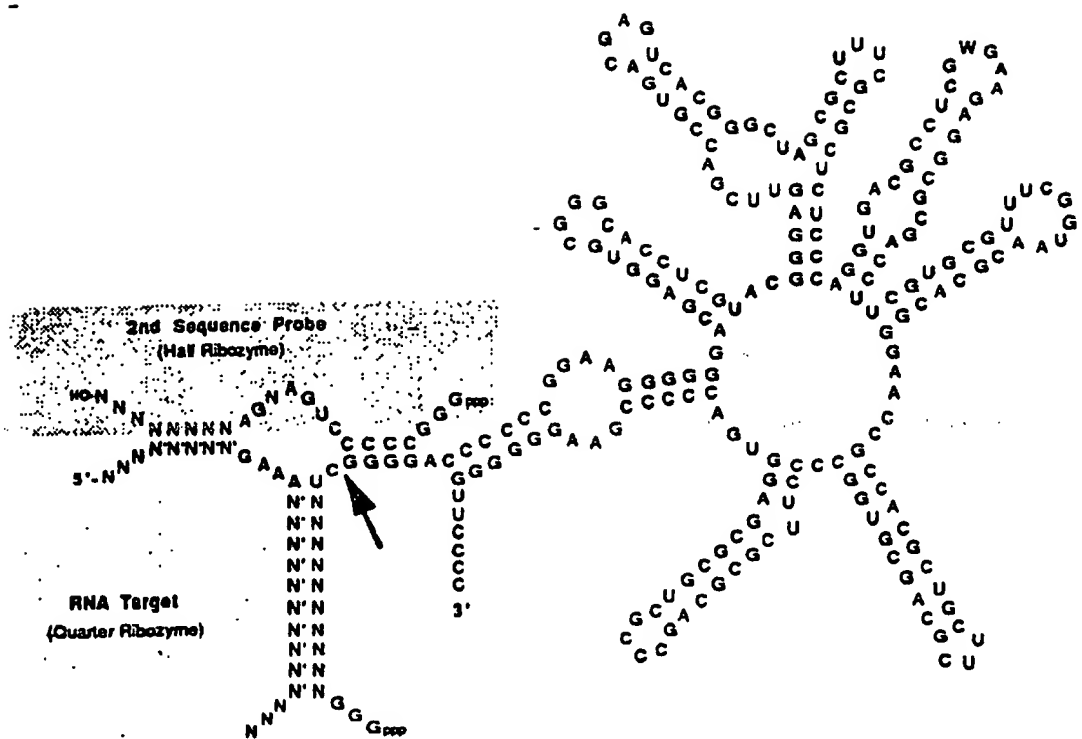


FIGURE 3

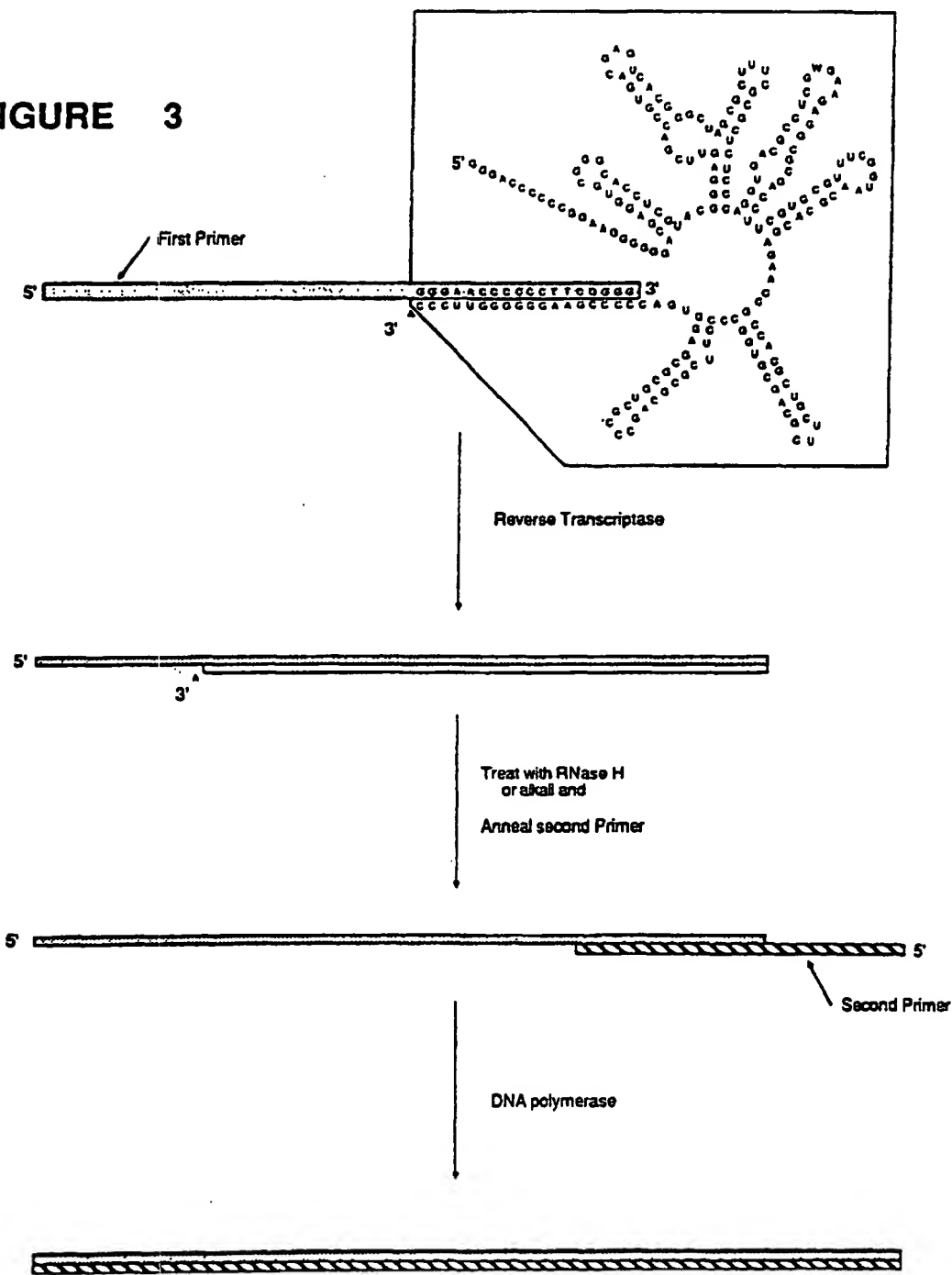


FIGURE 4A

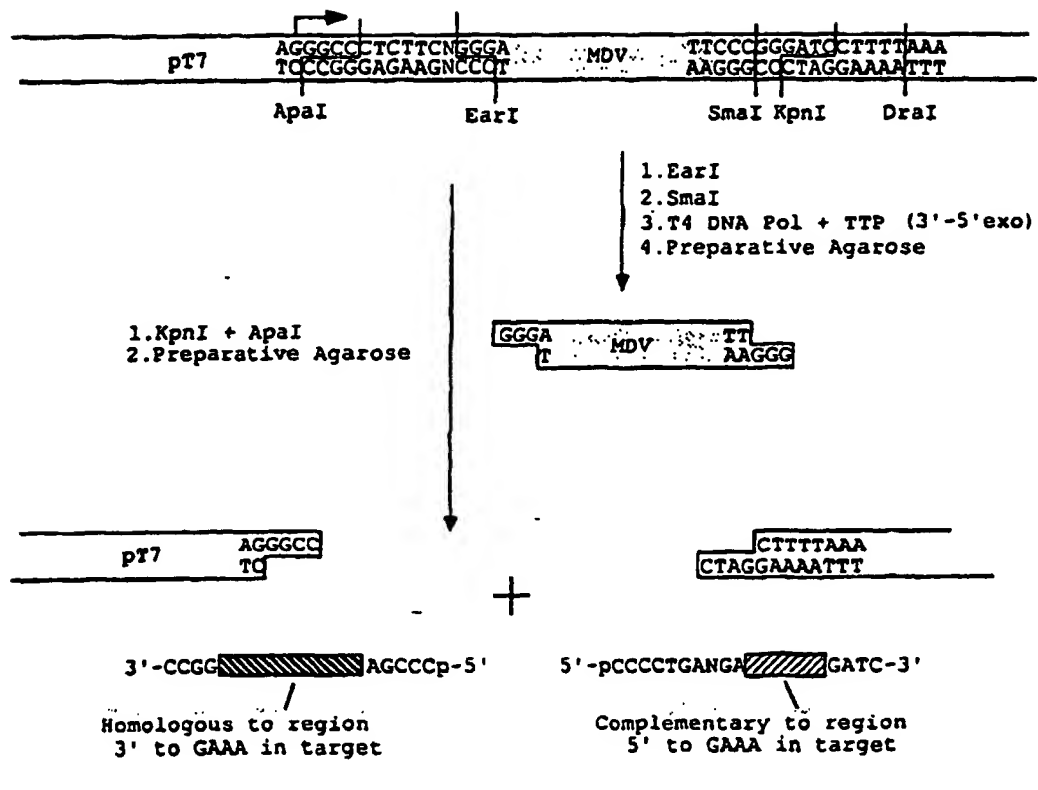


FIGURE
4B

FIGURE 4B

from Figure 4A

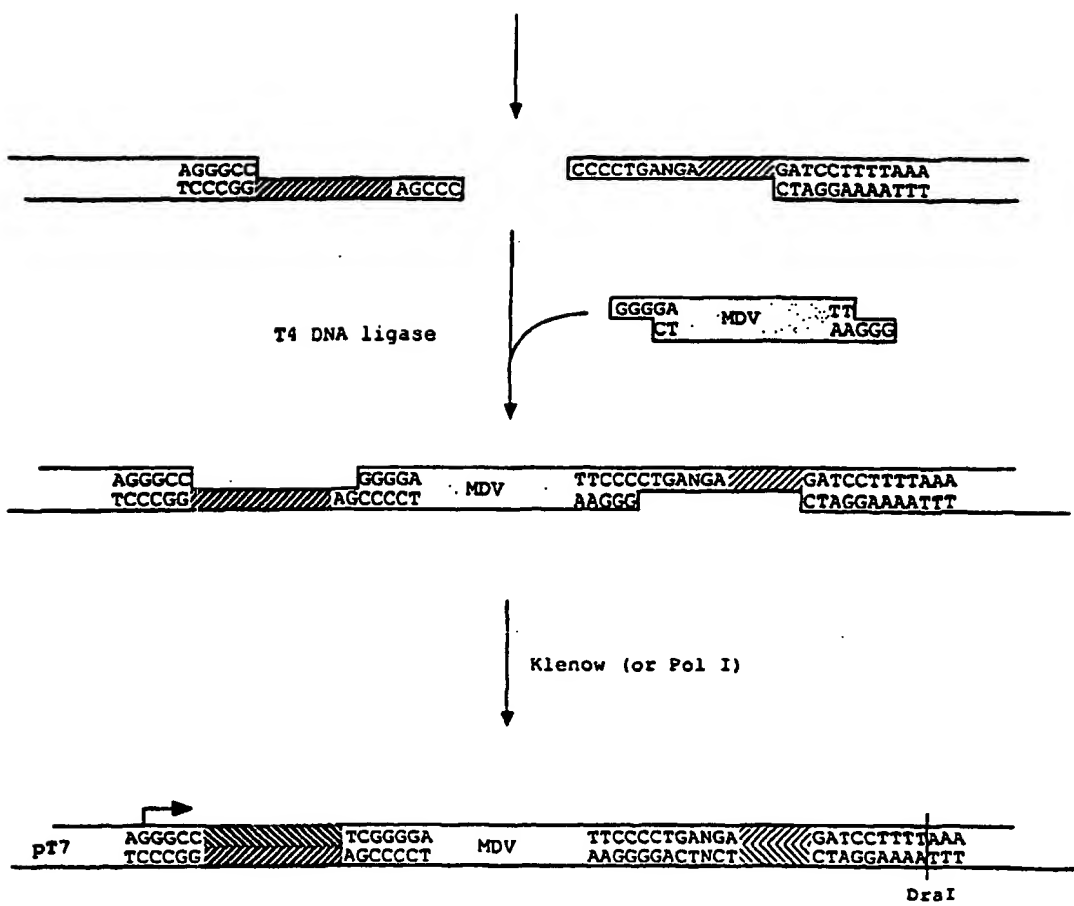


FIGURE 5A

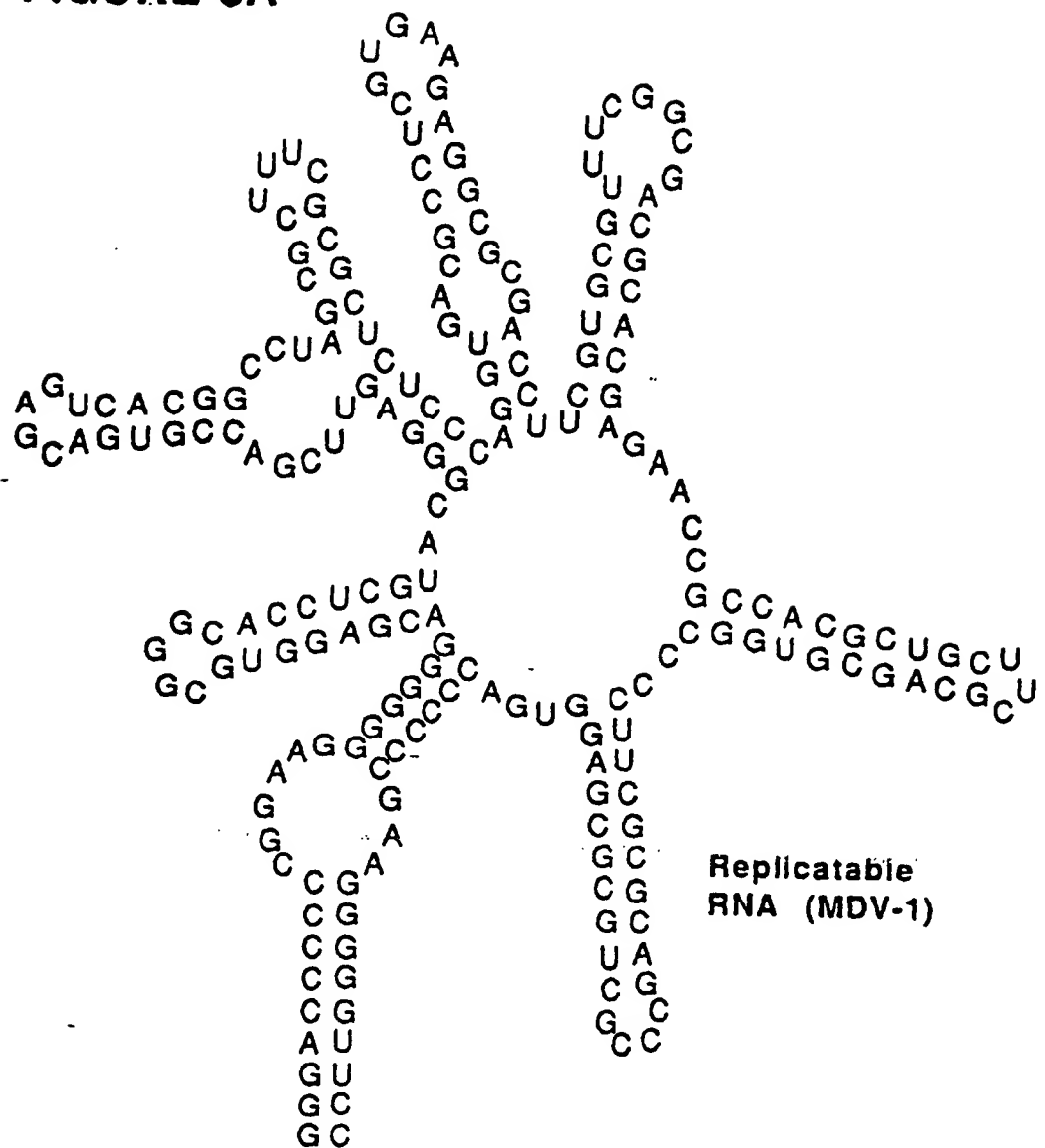


Figure 5B

FIGURE 5B

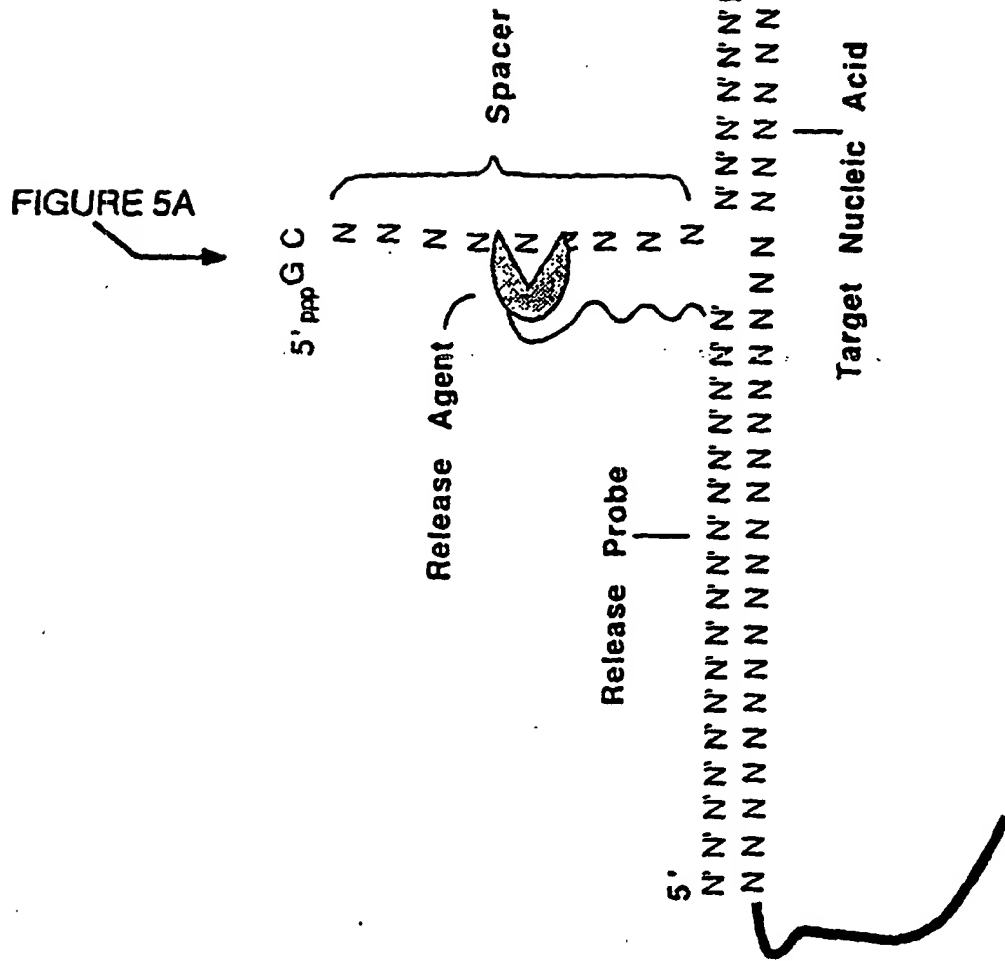


FIGURE 5A